CLAIMS

What is claimed is:

- 1. A method of producing an Isotope Map for data from a mass spectometric injection of a biological sample comprising:
 - a) performing Noise reduction and Centroiding with a Noise Reduction Module on said data from a mass spectometric injection of a biological sample, and
 - b) generating an Isotope Map from the noise reduced and centroided data, thereby producing an Isotope Map.
- 2. The method of claim 1, wherein said biological sample is comprised of unlabeled biomolecules.
- 3. The method of claim 1, wherein said biological sample is comprised of underivatized biomolecules.
- 4. The method of claim 1, wherein said biological sample is comprised of biomolecules that are both unlabeled and underivatized.
- 5. The method of claim 1, wherein said biological sample is comprised of cleaved biomolecules.
- 6. The method of claim 5, wherein said biomolecules are cleaved with an enzyme.
- 7. The method of claim 6, wherein said enzyme is trypsin.
- 8. A method for producing a Peptide Map from an Isotope Map for data from a

mass spectometric injection of a biological sample comprising:

- a) performing Peptide Detection on an Isotope Map with a Peptide Detection Module, and
- b) generating a Peptide Map from the results of the peptide detection, thereby producing a Peptide Map.
- 9. A method for determining the functions to align Peptide Maps from a pair of Peptide Maps, each representing a mass spectrometric injection of a biological sample, comprising:
 - a) performing Peptide Map Alignment on a pair of Peptide Maps with a Peptide Map Alignment Module, and
 - b) determining a column offset between the pair of injections, and
 - c) determining a retention time transformation function between the pair of injections.
- 10. A method for Alignment of Peptide Maps from a pair of Peptide Maps, each representing a mass spectrometric injection of a biological sample, comprising performing Peptide Map Alignment on a pair of Peptide Maps with a Peptide Map Alignment Module, and thereby producing Aligned Peptide Maps.
- 11. A method of determining differentially and uniquely expressed biomolecules from a pair of mass spectrometric injections of biological samples, comprising:
 - a) producing Isotope Maps for the samples;
 - b) producing Peptide Maps from the Isotope Maps;
 - c) aligning the Peptide Maps, and
 - d) determining the differences between the aligned maps.
- 12. A method of matching biomolecules from an LC-MS injection to LC-MS-MS fragmentation spectra from comprising:
 - a) producing Isotope Maps for the injections;
 - b) producing Peptide Maps from the Isotope Maps;
 - c) aligning the Peptide Maps, and
 - d) determining the matching biomolecules.

- 13. A computer implemented method of producing an Isotope Map for data from a mass spectometric injection of a biological sample comprising:
 - a) inputting data from a mass spectometric injection of a biological sample, and performing Noise reduction and Centroiding with a Noise Reduction Module on said, and
 - b) generating an Isotope Map from the noise reduced and centroided data, thereby producing an Isotope Map.
- 14. The computer implemented method of claim 13, wherein said biological sample is comprised of unlabeled biomolecules.
- 15. The computer implemented method of claim 13, wherein said biological sample is comprised of underivatized biomolecules.
- 16. The computer implemented method of claim 13, wherein said biological sample is comprised of biomolecules that are both unlabeled and underivatized.
- 17. The computer implemented method of claim 13, wherein said biological sample is comprised of cleaved biomolecules.
- 18. The computer implemented method of claim 17, wherein said biomolecules are cleaved with an enzyme.
- 19. The computer implemented method of claim 18, wherein said enzyme is trypsin.
- 20. A computer implemented method for producing a Peptide Map from an Isotope Map for data from a mass spectometric injection of a biological sample comprising:
 - a) inputting an Isotope Map;
 - b) performing Peptide Detection on said Isotope Map with a Peptide Detection Module, and

- c) generating a Peptide Map from the results of the peptide detection, thereby producing a Peptide Map.
- 21. A computer implemented method for determining the functions to align Peptide Maps from a pair of Peptide Maps, each representing a mass spectrometric injection of a biological sample, comprising:
 - a) inputting a pair of Peptide Maps;
 - b) performing Peptide Map Alignment on a pair of Peptide Maps with a Peptide Map Alignment Module;
 - c) determining a column offset between the pair of injections, and
 - d) determining a retention time transformation function between the pair of injections.
- 22. A computer implemented method for Alignment of Peptide Maps from a pair of Peptide Maps, each representing a mass spectrometric injection of a biological sample, comprising inputting a pair of Peptide Maps, performing Peptide Map Alignment on said pair of Peptide Maps with a Peptide Map Alignment Module, and thereby producing Aligned Peptide Maps.
- 23. A computer implemented method of determining differentially and uniquely expressed biomolecules from a pair of mass spectrometric injections of biological samples, comprising:
 - a) inputting data from said injections;
 - b) producing Isotope Maps for the samples;
 - c) producing Peptide Maps from the Isotope Maps;
 - d) aligning the Peptide Maps, and
 - e) determining the differences between the aligned maps.
- 24. A computer implemented method of matching biomolecules from an LC-MS injection to LC-MS-MS fragmentation spectra from comprising:
 - a) inputting data from said injections;
 - b) producing Isotope Maps for the injections;
 - c) producing Peptide Maps from the Isotope Maps;
 - d) aligning the Peptide Maps, and

- e) determining the matching biomolecules.
- 25. A computer-readable memory having stored thereon a program for producing an Isotope Map for data from a mass spectometric injection of a biological sample comprising:
 - a) computer code that receives as input data from a mass spectometric injection of a biological sample, and performing Noise reduction and Centroiding with a Noise Reduction Module on said, and
 - b) and computer code that generates an Isotope Map from the noise reduced and centroided data, thereby producing an Isotope Map.
- 26. The computer-readable memory of claim 25, wherein said biological sample is comprised of unlabeled biomolecules.
- 27. The computer-readable memory of claim 25, wherein said biological sample is comprised of underivatized biomolecules.
- 28. The computer-readable memory of claim 25, wherein said biological sample is comprised of biomolecules that are both unlabeled and underivatized.
- 29. The computer-readable memory of claim 25, wherein said biological sample is comprised of cleaved biomolecules.
- 30. The computer-readable memory of claim 29, wherein said biomolecules are cleaved with an enzyme.
- 31. The computer-readable memory of claim 30, wherein said enzyme is trypsin.
- 32. A computer-readable memory having stored thereon a program for producing a Peptide Map from an Isotope Map for data from a mass spectometric injection of a biological sample comprising:
 - a) computer code that receives as input an Isotope Map;

- b) computer code that performs Peptide Detection on said Isotope Map with a Peptide Detection Module, and
- c) computer code that generates a Peptide Map from the results of the peptide detection, thereby producing a Peptide Map.
- 33. A computer-readable memory having stored thereon a program for determining the functions to align Peptide Maps from a pair of Peptide Maps, each representing a mass spectrometric injection of a biological sample, comprising:
 - a) computer code that receives as input a pair of Peptide Maps;
 - b) computer code that performs Peptide Map Alignment on a pair of Peptide Maps with a Peptide Map Alignment Module;
 - c) computer code that determines a column offset between the pair of injections, and
 - d) computer code that determines a retention time transformation function between the pair of injections.
- 34. A computer-readable memory having stored thereon a program for Alignment of Peptide Maps from a pair of Peptide Maps, each representing a mass spectrometric injection of a biological sample, comprising computer code that receives as input a pair of Peptide Maps, and computer code that performs Peptide Map Alignment on said pair of Peptide Maps with a Peptide Map Alignment Module, and thereby produces Aligned Peptide Maps.
- 35. A computer-readable memory having stored thereon a program for determining differentially and uniquely expressed biomolecules from a pair of mass spectrometric injections of biological samples, comprising:
 - a) computer code that receives as input data from said injections;
 - b) computer code that produces Isotope Maps for the samples;
 - c) computer code that produces Peptide Maps from the Isotope Maps;
 - d) computer code that aligns the Peptide Maps, and
 - e) computer code that determines the differences between the aligned maps.
- 36. A computer-readable memory having stored thereon a program for matching biomolecules

from an LC-MS injection to LC-MS-MS fragmentation spectra from comprising:

- a) computer code that receives as input data from said injections;
- b) computer code that produces Isotope Maps for the injections;
- c) computer code that produces Peptide Maps from the Isotope Maps;
- d) computer code that aligns the Peptide Maps, and
- e) computer code that determines the matching biomolecules.
- 37. A computer system for producing an Isotope Map for data from a mass spectometric injection of a biological sample comprising a processor and a memory coupled to said processor, said memory encoding one or more programs, said one or more programs causing said processor to perform a method comprising:
 - a) computer code that receives as input data from a mass spectometric injection of a biological sample, and performing Noise reduction and Centroiding with a Noise Reduction Module on said, and
 - b) computer code that generates an Isotope Map from the noise reduced and centroided data, thereby producing an Isotope Map.
- 38. The computer system of claim 37, wherein said biological sample is comprised of unlabeled biomolecules.
- 39. The computer system of claim 37, wherein said biological sample is comprised of underivatized biomolecules.
- 40. The computer system of claim 37, wherein said biological sample is comprised of biomolecules that are both unlabeled and underivatized.
- 41. The computer system of claim 37, wherein said biological sample is comprised of cleaved biomolecules.
- 42. The computer system of claim 41, wherein said biomolecules are cleaved with an enzyme.

- 43. The computer system of claim 42, wherein said enzyme is trypsin.
- 44. A computer system for producing a Peptide Map from an Isotope Map for data from a mass spectometric injection of a biological sample comprising a processor and a memory coupled to said processor, said memory encoding one or more programs, said one or more programs causing said processor to perform a method comprising:
 - a) computer code that receives as input an Isotope Map;
 - b) computer code that performs Peptide Detection on said Isotope Map with a Peptide Detection Module, and
 - c) computer code that generates a Peptide Map from the results of the peptide detection, thereby producing a Peptide Map.
- 45. A computer system for determining the functions to align Peptide Maps from a pair of Peptide Maps, each representing a mass spectrometric injection of a biological sample, comprising a processor and a memory coupled to said processor, said memory encoding one or more programs, said one or more programs causing said processor to perform a method comprising:
 - a) computer code that receives as input a pair of Peptide Maps;
 - b) computer code that performs Peptide Map Alignment on a pair of Peptide Maps with a Peptide Map Alignment Module;
 - c) computer code that determines a column offset between the pair of injections, and
 - d) computer code that determines a retention time transformation function between the pair of injections.
- 46. A computer system for Alignment of Peptide Maps from a pair of Peptide Maps, each representing a mass spectrometric injection of a biological sample, comprising a processor and a memory coupled to said processor, said memory encoding one or more programs, said one or more programs causing said processor to perform a method comprising computer code that receives as input a pair of Peptide Maps, and computer code that performs Peptide Map Alignment on said pair of Peptide Maps with a Peptide Map Alignment Module, and thereby

produces Aligned Peptide Maps.

- 47. A computer system for determining differentially and uniquely expressed biomolecules from a pair of mass spectrometric injections of biological samples, comprising a processor and a memory coupled to said processor, said memory encoding one or more programs, said one or more programs causing said processor to perform a method comprising:
 - a) computer code that receives as input data from said injections;
 - b) computer code that produces Isotope Maps for the samples;
 - c) computer code that produces Peptide Maps from the Isotope Maps;
 - d) computer code that aligns the Peptide Maps, and
 - e) computer code that determines the differences between the aligned maps.
- 48. A computer system for matching biomolecules from an LC-MS injection to LC-MS-MS fragmentation spectra from comprising a processor and a memory coupled to said processor, said memory encoding one or more programs, said one or more programs causing said processor to perform a method comprising:
 - a) computer code that receives as input data from said injections;
 - b) computer code that produces Isotope Maps for the injections;
 - c) computer code that produces Peptide Maps from the Isotope Maps;
 - d) computer code that aligns the Peptide Maps, and
 - e) computer code that determines the matching biomolecules.
- 49. A method for displaying information on an Isotope Map for data from a mass spectometric injection of a biological sample comprising:
 - a) inputting data from a mass spectometric injection of a biological sample;
 - b) performing Noise reduction and Centroiding with a Noise Reduction Module on said data;
 - b) generating an Isotope Map from the noise reduced and centroided data, thereby producing an Isotope Map, and
 - c) displaying information on said Isotope Map to a user..

- 50. The method of claim 49, wherein said biological sample is comprised of unlabeled biomolecules.
- 51. The method of claim 49, wherein said biological sample is comprised of underivatized biomolecules.
- 52. The method of claim 49, wherein said biological sample is comprised of biomolecules that are both unlabeled and underivatized.
- 53. The method of claim 49, wherein said biological sample is comprised of cleaved biomolecules.
- 54. The method of claim 53, wherein said biomolecules are cleaved with an enzyme.
- 55. The method of claim 54, wherein said enzyme is trypsin.
- 56. A method for displaying information on a Peptide Map produced from an Isotope Map for data from a mass spectometric injection of a biological sample comprising:
 - a) inputting an Isotope Map;
 - b) performing Peptide Detection on said Isotope Map with a Peptide Detection Module;
 - c) generating a Peptide Map from the results of the peptide detection, thereby producing a Peptide Map, and
 - d) displaying information on said Peptide Map to a user.
- 57. A method for displaying information on the functions to align Peptide Maps from a pair of Peptide Maps, each representing a mass spectrometric injection of a biological sample, comprising:
 - a) inputting a pair of Peptide Maps;
 - b) performing Peptide Map Alignment on a pair of Peptide Maps with

- a Peptide Map Alignment Module;
- c) determining a column offset between the pair of injections;
- d) determining a retention time transformation function between the pair of injections, and
- e) displaying information on said functions to a user.
- 58. A method for displaying information on Alignment of Peptide Maps from a pair of Peptide Maps, each representing a mass spectrometric injection of a biological sample, comprising inputting a pair of Peptide Maps, and performing Peptide Map Alignment on said pair of Peptide Maps with a Peptide Map Alignment Module, thereby producing Aligned Peptide Maps, displaying information on said Aligned Peptide Maps to a user.
- 59. A method for displaying information on differentially and uniquely expressed biomolecules from a pair of mass spectrometric injections of biological samples, comprising:
 - a) inputting data from said injections;
 - b) producing Isotope Maps for the samples;
 - c) producing Peptide Maps from the Isotope Maps;
 - d) aligning the Peptide Maps;
 - e) determining the differences between the aligned maps, and
 - f) displaying information on said differences to a user.
- 60. A method for displaying information on matching biomolecules from an LC-MS injection to LC-MS-MS fragmentation spectra from comprising:
 - a) inputting data from said injections;
 - b) producing Isotope Maps for the injections;
 - c) producing Peptide Maps from the Isotope Maps;
 - d) aligning the Peptide Maps, and
 - e) determining the matching biomolecules,
 - f) and displaying information on said matching biomolecules to a user.